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L3 L4 L5 L6	FILE	'HCAPLUS' ENTERED AT 13:08:44 ON 18 SEP 2008 1990 S (L1/THU) OR (L2/THU) 37844 S CYCLODEXTRIN 47 S L3 AND L4 15 S L5 AND (PY<2003 OR AY<2003 OR PRY<2003)
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=> file hcaplus
COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

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FILE COVERS 1907 - 18 Sep 2008 VOL 149 ISS 12 FILE LAST UPDATED: 16 Sep 2008 (20080916/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> file registry
COST IN U.S. DOLLARS

SINCE FILE TOTAL
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FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 16 SEP 2008 HIGHEST RN 1049663-83-3 DICTIONARY FILE UPDATES: 16 SEP 2008 HIGHEST RN 1049663-83-3

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TSCA INFORMATION NOW CURRENT THROUGH July 5, 2008.

Please note that search-term pricing does apply when conducting ${\tt SmartSELECT}$ searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

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E2
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E.3
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ARACHIDONO/BI
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COST IN U.S. DOLLARS
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FULL ESTIMATED COST 10.76 13.66

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FILE COVERS 1907 - 18 Sep 2008 VOL 149 ISS 12 FILE LAST UPDATED: 16 Sep 2008 (20080916/ED)

HCAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2008.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

 \Rightarrow s (11/thu) or (12/thu) 33873 L1 1048155 THU/RL 742 L1/THU (L1 (L) THU/RL) 41765 L2 1048155 THU/RL 1604 L2/THU (L2 (L) THU/RL) L3 1990 (L1/THU) OR (L2/THU) => s cyclodextrin L437844 CYCLODEXTRIN => s 13 and 14 47 L3 AND L4 T.5 => s 15 and (PY<2003 or AY<2003 or PRY<2003) 22958910 PY<2003 4497131 AY<2003 3965546 PRY<2003 L6 15 L5 AND (PY<2003 OR AY<2003 OR PRY<2003) => d 16 1-15 ti abs bib ANSWER 1 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN 1.6 ΤI Antler composition containing a matrix comprising β cyclodextrin, an ester, and proteinase inhibitor

AB An antler composition and its manufacturing process are disclosed, which comprises an

antler extract mixture and a matrix which comprises β - cyclodextrin , a higher ester compound, a proteinase inhibitor, and an organic solvent;

wherein the weight ratio of the matrix to the antler extract mixture is between 1:1.5 and 1:2.7. The antler composition poses excellent activities and stable properties to be released steadily in human body. The present invention also relates to the antler extract mixture and the process for preparing the antler composition and the antler extract mixture 2003:971301 HCAPLUS <<LOGINID::20080918>> ΑN DN140:19864 ΤI Antler composition containing a matrix comprising β cyclodextrin, an ester, and proteinase inhibitor Hsu, David H.; Chen, Eve Sze-Ju INPAU.S. Pat. Appl. Publ., 10 pp. SO CODEN: USXXCO DTPatent English LAFAN.CNT 1 PATENT NO. KIND APPLICATION NO. DATE DATE -----____ _____ _____ US 20030228372 A1 20031211 US 2002-325746 20021223 <--PΙ US 7005144 В2 20060228 PRAI TW 2002-91112440 Α 20020607 <--TW 2002-91112441 Α 20020607 <--20020607 <--TW 2002-91112442 Α RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 2 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN L6 ΤI Topical formulations of resorcinols and cannabinoids and methods of use AΒ The invention provides a method for preventing the transmission of HIV from one individual to another. In accordance with the method, a pharmacol. acceptable composition including at least one resorcinol derivative and/or cannabinoid (e.g., cannabinol derivs., $\Delta 8-THC$ derivs., cannabichromene derivs., cannabidiol derivs., cannabigerol derivs.) (including combinations thereof) is administered topically to a first individual harboring HIV, or to a second individual at risk of infection with HIV, proximate in time with contact between the first individual and the second individual. The invention also provides topical formulations of at least one resorcinol and/or cannabinoid and water insol. polymers as 2003:777578 HCAPLUS <<LOGINID::20080918>> ΑN DN 139:296973 ΤI Topical formulations of resorcinols and cannabinoids and methods of use ΤN Travis, Craiq R. PAImmugen Pharmaceuticals, Inc., USA PCT Int. Appl., 68 pp. SO CODEN: PIXXD2 DTPatent LAEnglish FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE ____ _____ ______ WO 2003080043 A1 20031002 WO 2003-US8314 PI20030318 <--W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,

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A1 20031002 WO 2003-US8314 20030318 <---
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RE.CNT 1
              THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
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- L6 ANSWER 3 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Quality study of volatile oil enclosed with $\beta\text{--}$ cyclodextrin in Naokangling capsule
- AB Study the quality of volatile oil enclosed with β -cyclodextrin (β -CD) in Naokangling capsule. The quality of the volatile oil in Naokangling capsule before and after enclosure was examined by thin layer chromatog., UV and gas chromatog.-mass spectrometry. The inclusion of volatile oil and β -cyclodextrin was steady, and the quality of volatile oil was not changed before and after enclosure. The process of enclosure with β -CD can keep the active components of the volatile oil in Naokangling capsule.
- AN 2003:554493 HCAPLUS <<LOGINID::20080918>>
- DN 140:258803
- TI Quality study of volatile oil enclosed with $\beta-$ cyclodextrin in Naokangling capsule
- AU Wang, Yan; Zhou, Liling; Liu, Qingfei; Qiu, Meixian; Liang, Shuyan
- CS Guangzhou University of TCM, Canton, 510405, Peop. Rep. China
- SO Guangzhou Zhongyiyao Daxue Xuebao (2002), 19(4), 311-313 CODEN: GZDXFQ; ISSN: 1007-3213
- PB Guangzhou Zhongyiyao Daxue Xuebao Bianjibu
- DT Journal
- LA Chinese
- L6 ANSWER 4 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- ${\tt TI}$ Compositions comprising an o/w emulsion containing conjugated linoleic acid
- AB The present invention provides a method of treatment of a human or non-human (e.g. mammalian, avian or reptilian) animal subject by the parenteral administration of a lipophilic pharmaceutical agent, the improvement comprising administering said pharmaceutical agent in an oil-in-water emulsion containing a conjugated linoleic acid (CLA) or a physiol. tolerable derivative thereof. A mixture of 10 g CLA triglyceride (produced by reacting CLA with glycerol), 1.0 g purified egg phospholipid, 50 mg sodium stearate and 5 g α -tocopherol was finely dispersed. A mixture of 100 mL water containing 2.5 g glycerol and 0.05 mmol NaOH was added to the CLA mixture during stirring at room temperature. The mixture was homogenized
 - in a high pressure homogenator and the final emulsion filled into vials and heat-sterilized.
- AN 2002:695819 HCAPLUS <<LOGINID::20080918>>
- DN 137:222086
- ${\tt TI}$ Compositions comprising an o/w emulsion containing conjugated linoleic acid
- IN Remmereit, Jan; Klaveness, Jo
- PA Natural Asa, Norway; Cockbain, Julian
- SO PCT Int. Appl., 29 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 1

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PRAI GB 2001-5622
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                              20020307 <--
    WO 2002-GB996
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             THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 15
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ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN L6

Cubic liquid crystalline compositions and methods for their preparation ΤI

A dry powder cubic gel precursor comprising an encapsulating compound, an amphiphile capable of forming a cubic liquid crystalline phase, and optionally

solvent is described. The encapsulating compound (A), amphiphile (B), and optional solvent (C) are present in mass fractions relative to each other such that 1.0 = a + b + c, wherein a is the mass fraction of A, b is the mass fraction of B, and c is the mass fraction of C. Further, 1.0 > a >0, 1.0 > b > 0, 1.0 > c > 0 and a, b, and c do not fall within a cubic liquid crystalline phase region on a phase diagram representing phase behavior

of

а

A, B, and C. A method of making the cubic gel precursor comprises the steps of: (i) dissolving an encapsulating compound in a solvent; (ii) adding an amphiphile; (iii) mixing the encapsulating compound and amphiphile, wherein steps (i), (ii), and (iii) are performed in any order; (iv) atomizing the mixture obtained; and, (v) drying the mixture For example, an active ingredient (fatty acid solution) was encapsulated in powders made by spray-drying a liquid solution The liquid solution was prepared from a premix

water and 33% starch at 70° . A second solution of 90% monoolein and 10% fatty acid mix (20% omega-3, 80% triglyceride oil) was prepared at 60° . The oil solution was then added to the starch-water solution forming a 9% monoolein, 30% starch, 60% water, and 1% fatty acid mixture A high shear mixing system was used to keep the system mixed and maintained above 90°. The mixture was then pumped at a rate of 8 mL/min through the liquid side of a twin-fluid atomizer, with slight adjustments being made to the flow rate to keep the temperature of the exit air in the system between 90-100°. The liquid feed was atomized with air at a pressure of 42.6 psi (293.5 kPa). Upon drying, the powder has a composition of 22.5% monoolein, 75% starch, and 2.5% fatty acid mixture The powder appears to exhibit a bimodal size distribution of larger 10 μm particles and smaller 3-5 μm particles, all of which exhibit the classical shrinkage that is characteristic of starch capsules during their cooling. The uniform appearance of the powders can be an excellent indicator that the fatty acid active is encapsulated within the starch shells.

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137:206536
DN
ΤI
     Cubic liquid crystalline compositions and methods for their preparation
     Spicer, Patrick Thomas; Small, William Broderick, II; Lynch, Matthew
IN
     Lawrence
     The Procter & Gamble Company, USA
PA
SO
     PCT Int. Appl., 37 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
                        KIND DATE
                                                APPLICATION NO.
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     WO 2002066014 A2 20020829
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EP 1361865 A2 20031119 EP 2002-721031
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WO 2002-US4776 W 20020219 <--
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L6
     ANSWER 6 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
ΤI
     Skin sanitizing compositions
AΒ
     The present invention relates to compns. and methods of sanitizing and
     moisturizing skin surfaces. A sanitizing and moisturizing gel contained
     EtOH 55, isopropanol 3, Biowax-754 0.4, Carbopol Ultrez-10 0.3, Carbowax
     PEG-200 0.26, propylene glycol 0.02, aminomethylpropanol 0.15, and perfume
     0.1%, and water qs.
     2002:551533 HCAPLUS <<LOGINID::20080918>>
AN
DN
     137:114518
     Skin sanitizing compositions
TI
     Sine, Mark Richard; Wei, Karl Shiqing; Jakubovic, David Andrew; Thomas,
IN
     Cheyne P.; Dodd, Michael Thomas; Putman, Christopher Dean
PA
     The Procter & Gamble Company, USA
     U.S., 14 pp., Cont. of U.S. Ser. No. 321,291.
     CODEN: USXXAM
DT
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LA
     English
FAN.CNT 2
     PATENT NO.
                          KIND DATE APPLICATION NO.
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PT
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2002:657934 HCAPLUS <<LOGINID::20080918>>

AN

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US 1999-120098P P 19990216 <--

US 1999-321291 A2 19990527 <--
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RE.CNT 56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 7 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Clear oil-containing pharmaceutical compositions containing a therapeutic agent
- AB The present invention relates to pharmaceutical compns. and methods for improved solubilization of triglycerides and improved delivery of therapeutic agents. Compns. of the present invention include a carrier, where the carrier is formed from a combination of a triglyceride and at least 2 surfactants, at least one of which is hydrophilic. Upon dilution with an aqueous medium, the carrier forms a clear, aqueous dispersion of the triglyceride and surfactants. Thus, a formulation contained soybean oil, 80, Tween-20 200, and Tween-80 800 mg.
- AN 2002:185694 HCAPLUS <<LOGINID::20080918>>
- DN 136:252483
- TI Clear oil-containing pharmaceutical compositions containing a therapeutic agent
- IN Chen, Feng-Jing; Patel, Mahesh V.; Fikstad, David T.
- PA Lipocine, Inc., USA
- SO U.S. Pat. Appl. Publ., 45 pp., Cont.-in-part of U.S. Ser. No. 751,968. CODEN: USXXCO
- DT Patent
- LA English

FAN.CNT 13

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	US	20010024658	A1	20010927	US	2000-751968	20001229	<
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RE.CNT 88 THERE ARE 88 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 8 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- ${\tt TI}$ Triglyceride-free compositions and methods for enhanced absorption of hydrophilic therapeutic agents
- AB The present invention relates to triglyceride-free pharmaceutical compns., pharmaceutical systems, and methods for enhanced absorption of hydrophilic therapeutic agents. The compns. and systems include an absorption enhancing carrier, where the carrier is formed from a combination of at

least two surfactants, at least one of which is hydrophilic. A hydrophilic therapeutic agent can be incorporated into the composition, or can be co-administered with the composition as part of a pharmaceutical system. The invention also provides methods of treatment with hydrophilic therapeutic agents using these compns. and systems. For example, when a composition containing Cremophor RH40 0.30, Arlacel 186 0.20, Na taurocholate 0.18,

and propylene glycol 0.32 g, resp., was used, the relative absorption of PEG 4000 as a model macromol. drug was enhanced by 991%.

- AN 2001:136991 HCAPLUS <<LOGINID::20080918>>
- DN 134:198075
- TI Triglyceride-free compositions and methods for enhanced absorption of hydrophilic therapeutic agents
- IN Patel, Mahesh V.; Chen, Feng-Jing
- PA Lipocine, Inc., USA
- SO PCT Int. Appl., 113 pp.

CODEN: PIXXD2

- DT Patent
- LA English

FAN.CNT 13

11111	PATENT NO.					KIND DATE			APPLICATION NO.						DATE			
ΡI	WO	20010	121	55		A1		2001	0222		WO 2	000-1	US18	807		2	0000	710 <
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			SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	UZ,	VN,	YU,
			ZA,	ZW														
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	EР																	710 <
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																		710 <
		7808																710 < 710 <
		20010																229 <
		64583				B2		2001			05 2	000-	1313	00		۷.	3001	229 \
PRAT		1999-						1999			_							
		2000-						2000										
RE.C	_	1							-			ABLE	FOR	THI	S RE	CORD		

- L6 ANSWER 9 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Clear aqueous dispersions of triglycerides and surfactants for delivery of drugs and nutrients

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- AB The present invention relates to drug and nutrient delivery systems, and in particular to pharmaceutical compns. and methods for improved solubilization of triglycerides and improved delivery of therapeutic agents. Compns. of the present invention include a triglyceride and a carrier, where the carrier is formed from a combination of at least two surfactants, at least one of which is hydrophilic. Upon dilution with an aqueous
- solvent, the composition forms a clear, aqueous dispersion of the triglyceride and
 - surfactants. An optional therapeutic agent can be incorporated into the composition, or can be co-administered with the composition. The invention also

provides methods of enhancing triglyceride solubility and methods of treatment with therapeutic agents using these compns. Several formulations were presented of compns. that can be prepared according to the present invention using a variety of therapeutic agents. Examples of aqueous dispersions include: (1) Cremophor RH-40 0.75, Peceol 0.25, corn oil 0.40, and fenofibrate 0.10; (2) Cremophor RH-40 0.57, Crovol M-40 0.43, corn oil 0.40, and Rofecoxib 0.15; (3) Tween 80 0.70, Tween 85 0.35, Miglyol 812 0.30, Paclitaxel 0.10, and PEG 400 0.25; or (4) Kessco PEG 400 MO 0.33, corn oil 0.30, and Terbinafine 0.25 parts, resp.

- AN 2001:31306 HCAPLUS <<LOGINID::20080918>>
- DN 134:105846
- TI Clear aqueous dispersions of triglycerides and surfactants for delivery of drugs and nutrients
- IN Chen, Feng-Jing; Patel, Mahesh V.
- PA Lipocine, Inc., USA
- SO PCT Int. Appl., 103 pp.

CODEN: PIXXD2

- DT Patent
- LA English

FAN.CNT 13

	PATENT NO. KIND)	DATE		1	APPL	ICAT	ION I	NO.		DATE					
ΡI	WO	2001	0019	 60		A1	_	2001	0111	Ţ	WO 2	000-1	JS15:	133		2	0000	602	<
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PRAI	US	1999	-3450					1999	0630	<	-								
	MO	2000	-US1	5133		\mathbb{W}		2000	0602	<	-								
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- L6 ANSWER 10 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Pharmaceutical compositions and methods for improved delivery of hydrophobic therapeutic agents

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- AB The present invention relates to triglyceride-free pharmaceutical compns. for delivery of hydrophobic therapeutic agents. Compns. of the present invention include a hydrophobic therapeutic agent and a carrier, where the carrier is formed from a combination of a hydrophilic surfactant and a hydrophobic surfactant. Upon dilution with an aqueous solvent, the composition forms
- a clear, aqueous dispersion of the surfactants containing the therapeutic agent.

The invention also provides methods of treatment with hydrophobic therapeutic agents using these compns. A pharmaceutical composition contained cyclosporin 0.14, Cremophor RH-40 0.41, Arlacel186 0.29, sodium taurocholate 0.26, and propylene glycol 0.46 mg.

- AN 2000:608551 HCAPLUS <<LOGINID::20080918>>
- DN 133:213151

```
hydrophobic therapeutic agents
     Patel, Manesh V.; Chen, Feng-Jing
ΙN
     Lipocine, Inc., USA
PA
SO
     PCT Int. Appl., 98 pp.
     CODEN: PIXXD2
DT
     Patent
     English
FAN.CNT 13
     PATENT NO.
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    WO 2000050007 A1 20000831 WO 2000-US165 20000105 <--
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                      B1 20010925 US 1999-258654
A1 20000831 CA 2000-2365536
A 20000914 AU 2000-22242
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                                                                  19990226 <--
                        A1
A
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NZ 2000-513810
     JP 2002537317 T
                        A 20040227 N:
A 19990226 <--
                            20021105
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     NZ 513810
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PRAI US 1999-258654
     WO 2000-US165
             THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 4
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 11 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
L6
ΤI
    Pharmaceutical composition with base of coconut oil and its use
     A composition containing cetyl alc., coconut oil, polyoxyethylene oleo-linoleic
AΒ
     glyceride, optionally water, and optionally other additives and/or
     pharmaceutically active principle(s), said composition containing in % of dry
     matter: 5 to 15 % of polyoxyethylene oleo-linoleic glyceride, and 20 to 40
     % of cetyl alc., the ratio by weight of coconut oil and other additives
     and/or pharmaceutically active principle(s)/cetyl alc. ranging between 2/1
     and 80/15. A powder contained coconut oil 60, cetyl alc. 25, labrafil 5,
     and essential oils 5 parts. The powder is mixed with an equal amts. of
     water to make a cream.
    1998:568741 HCAPLUS <<LOGINID::20080918>>
ΑN
DN 129:180166
OREF 129:36509a,36512a
ΤI
    Pharmaceutical composition with base of coconut oil and its use
IN
     Streels, Elisabeth
PA
SO
     PCT Int. Appl., 20 pp.
     CODEN: PIXXD2
DT
    Patent
    French
LA
FAN.CNT 1
                       KIND DATE APPLICATION NO.
     PATENT NO.
                                                                 DATE
                A1 19980820 WO 1997-BE16 19970214 <--
    WO 9835700
РΤ
        W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC,
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Pharmaceutical compositions and methods for improved delivery of

ТΙ

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LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT,
          RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR,
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     AU 9717611
                                    19980908
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                             Α
                                                                           19970214 <--
     EP 957938
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     CA 2383806
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                                                                           20010420 <--
PRAI WO 1997-BE16
                                    19970214
                             Α
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     WO 2001-BE70
                             TAT
                                    20010420
                                              <--
               THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
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- RE.CNT 3 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- ANSWER 12 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN L6
- Percutaneous absorption and histopathology of a poloxamer-based ΤI formulation of capsaicin analog
- A new synthetic capsaicin analog (CA) modified with 4-hydroxyl and alkyl ABchain of capsaicin was synthesized as a potent anti-inflammatory analgesic drug and is now on clin. trial in Korea. The purpose of this study was to investigate the percutaneous absorption and histopathol. of a poloxamer-based formulation of CA. A poloxamer-based gel was prepared by cold method using poloxamer 407. Vertical Franz type diffusion cells were used for skin penetration of drug against receptor phase filled with about 10 mL of 0.9 isotonic saline at 32°C . The concentration of drug was determined by the reverse phased HPLC (C18, Symmetry®) with fluorometeric detector. Total amount of CA free base permeated was higher than that of the CA salt form. Percutaneous absorption of CA was greatly enhanced in ethanol and PG than that in water, 2-hydroxypropy- β cyclodextrin and PEG400. As ethanol concentration increased, percutaneous absorption greatly increased. The flux rate of CA increased slightly when PG was added to ethanol solution The marked enhancing effect of the 5 fatty acid IPM in cosolvents was also noted on the percutaneous absorption of a poloxamer-based formulation of CA. Addition of 5 OA and 5 LA into the gel containing 5 IPM resulted in a slight increase in skin permeation. No significant difference in skin permeation was observed as a function of poloxamer content (20, 25 and 30). The buffer system of 30 poloxamer-based gel slightly changed the cumulative amts. of CA penetrated for 24 h. The flux of poloxamer-based gels increased linearly as the drug concentration increased. There was a variation of percutaneous absorption of

drug, depending on the species used. The flux of a poloxamer-based formulation of CA was the highest in case of hairless mice but the lowest in hamsters. No skin erythema and histopathol. changes were observed on the dorsal site of hairless mice in six groups after a week or two months application, suggesting no skin toxicity of the poloxamer-based gel. Based on these findings, the current poloxamer-based formulation appears useful in the systemic delivery of CA as topical or transdermal patch formulations.

1997:790366 HCAPLUS <<LOGINID::20080918>> ΑN

128:93107 DN

the

OREF 128:18121a, 18124a

- Percutaneous absorption and histopathology of a poloxamer-based ΤI formulation of capsaicin analog
- ΑU Lee, Beom-Jin; Lee, Tae-Sup; Cha, Bong-Jin; Kim, Soon-Hoe; Kim, Won-Bae
- CS College of Pharmacy, Biological Rhythm and Controlled Release Laboratory, Kangwon National University, Chuncheon, 200-701, S. Korea
- SO International Journal of Pharmaceutics (1997), 159(1), 105-114 CODEN: IJPHDE; ISSN: 0378-5173

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PB Elsevier Science B.V.
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- DT Journal
- LA English
- RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L6 ANSWER 13 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Mucosal preparation containing physiologically active peptide
- AB This invention related to a mucosal preparation obtained by blending a physiol. active peptide at least with a sorbefacient and a vasodilatory compound Owing to the combined use of the sorbefacient with the vasodilatory compound, the absorption of any desired physiol. active peptide can be enhanced and thus it can be self-administered to a patient without giving any pain caused by parenteral injection. Therefore, it is highly useful as a preparation of a physiol. active peptide for prolonged administration. As the physiol. active peptide, use can be made of insulin, calcitonin, human PTH, somatostatin, glucagon, etc. As the sorbefacient, use can be made of bile acid salts, cyclodextrin, phospholipids, nonionic surfactants, higher fatty acids, etc. As the vasodilatory compds., use can be made of calcium channel inhibitors, prostaglandin E1, isosorbide nitrate, nitroglycerin, etc.
- AN 1997:259764 HCAPLUS <<LOGINID::20080918>>
- DN 126:242891
- OREF 126:46901a,46904a
- TI Mucosal preparation containing physiologically active peptide
- IN Yamamoto, Nakayuki; Ito, Teruomi
- PA Asahi Kasei Kogyo Kabushiki Kaisha, Japan; Hisamitsu Seiyaku Kabushiki Kaisha; Yamamoto, Nakayuki; Ito, Teruomi
- SO PCT Int. Appl., 48 pp. CODEN: PIXXD2
- DT Patent
- LA Japanese
- FAN.CNT 1

1 2 2 1 1 4 1	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 9706813	A1		WO 1996-JP2277	19960812 <
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	JP 11292787	A	19991026	JP 1995-208010	19950815 <
	CN 1179723	A	19980422	CN 1996-192821	19960812 <
	EP 845265	A1	19980603	EP 1996-926626	19960812 <
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	IE, FI				
	JP 3824023	B2	20060920	JP 1997-509140	19960812 <
PRAI	JP 1995-208010	A	19950815	<	
	WO 1996-JP2277	W	19960812	<	
OS	MARPAT 126:242891	L			

- L6 ANSWER 14 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Effects of topical anandamides on intraocular pressure in normotensive rabbits
- AB A series of anandamide-type compds. were synthesized and studied for their effect on the intraocular pressure (IOP) of normotensive pigmented rabbits. Each test compound was dissolved in an aqueous 2-hydroxypropyl- β -cyclodextrin solution and administered (31.25-62.5 μg) unilaterally to the eye. The most promising anandamides caused a statistically significant reduction of IOP in treated eyes, usually preceded by an initial transient elevation of IOP, compared to saline controls. In the contralateral untreated eyes, only a marginal or short hypotensive response was observed Indomethacin pre-treatment (12.5 mg, s.c.) eliminated the IOP response to administered anandamides and arachidonic acid,

```
indicating the involvement of prostaglandins. Structure-activity
    relations are discussed.
    1996:269518 HCAPLUS <<LOGINID::20080918>>
ΑN
    125:332
DN
OREF 125:55a,58a
    Effects of topical anandamides on intraocular pressure in normotensive
    rabbits
ΑU
    Pate, David W.; Jarvinen, Kristiina; Urtti, Arto; Jarho, Pekka; Fich,
    Mette; Mahadevan, Vaidyanath; Jarvinen, Tomi
CS
    Department Pharmaceitucal Chemistry, University Kuopio, Finland
    Life Sciences (1996), 58(21), 1849-60
SO
    CODEN: LIFSAK; ISSN: 0024-3205
PB
    Elsevier
DT
    Journal
LA
    English
    ANSWER 15 OF 15 HCAPLUS COPYRIGHT 2008 ACS on STN
1.6
    Transparent liquid for encapsulated drug delivery
TI
AΒ
    A stable transparent multi-component composition useful for the delivery of
    water soluble active agents to animals is provided. The compns. are
    formulated with a mixture of an oil phase, an aqueous phase, and a surfactant
    system, along with the active agent to be delivered to the animal. The
    compns. are specially formulated to be compatible with capsules such as
    gelatin and starch capsules. The aqueous phase of the compns. contains a
    substantial amount of polyethylene glycol and can optionally also contain a
    plasticizer. Preferred active agents are proteinaceous materials.
    Calcein bioavailability from a transparent liquid containing Captex 200 12,
    Imwitor 308 29.8, Tween 80 19.2, PEG 400 32.4, sorbitol 1.6, water 3%
    weight/weight, and 100 mM calcein solution in 10 mM Tris pH 7.4 3%
weight/weight, resp.,
    was studied.
    AN
DN
    123:152922
OREF 123:27049a,27052a
    Transparent liquid for encapsulated drug delivery
ΙN
    Yiv, Seang H.
PA
    Ibah, Inc., USA
SO
    PCT Int. Appl., 66 pp.
    CODEN: PIXXD2
DT
    Patent
LA
    English
FAN.CNT 2
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    PATENT NO.
                       KIND
                               DATE
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    WO 9514037
                       A1 19950526 WO 1994-US13394 19941116 <--
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            MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SI, SK, TJ, TT, UA,
            US, UZ
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            TD, TG
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    AU 692506
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    EP 736041
                                         EP 1995-904099
                                                                19941116 <--
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                              20060208
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    JP 09510182 T 19971014 JP 1994-514649 19941116 <--
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    US 5707648
                       A 19980113
                                          US 1995-406935
                                                                19950517 <--
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FILE 'HCAPLUS' ENTERED AT 13:08:00 ON 18 SEP 2008

FILE 'REGISTRY' ENTERED AT 13:08:12 ON 18 SEP 2008 EXP ARACHIDONIC

EXP ARACHIDONIC/CN

L1 1 S E5

EXP LINOLEIC/CN

L2 1 S E4

FILE 'HCAPLUS' ENTERED AT 13:08:44 ON 18 SEP 2008

L3 1990 S (L1/THU) OR (L2/THU)

L4 37844 S CYCLODEXTRIN

L5 47 S L3 AND L4

L6 15 S L5 AND (PY<2003 OR AY<2003 OR PRY<2003)

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COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 49.03 62.69 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -12.00-12.00

SESSION WILL BE HELD FOR 120 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 13:09:48 ON 18 SEP 2008

Connecting via Winsock to STN

Welcome to STN International! Enter x:X

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* * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * SESSION RESUMED IN FILE 'HCAPLUS' AT 13:17:16 ON 18 SEP 2008 FILE 'HCAPLUS' ENTERED AT 13:17:16 ON 18 SEP 2008 COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS) f

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CA SUBSCRIBER PRICE	-12.00	-12.00

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FULL ESTIMATED COST	49.03	62.69
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	ENTRY	SESSION
CA SUBSCRIBER PRICE	-12.00	-12.00

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STRUCTURE FILE UPDATES: 16 SEP 2008 HIGHEST RN 1049663-83-3 DICTIONARY FILE UPDATES: 16 SEP 2008 HIGHEST RN 1049663-83-3

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH July 5, 2008.

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=> exp alpha-	cyclodext/cn		
E1	1 ALPHA-CRYSTALI CN	LIN (GALLERIA MELLONELLA (GENE ALPHA-CRYS/SHSP)/
E2	•••	LIN B CHAIN (RATTUS NORVE	GICUS GENE ALPHA(B)-CR
E3	0> ALPHA-CYCLODE	XT/CN	
E 4	1 ALPHA-CYCLODEX 6)/CN	XTRINASE (GEOBACILLUS KAUS	STOPHILUS STRAIN HTA42
E5	• •	XYL-ALPHA-PHENYL-1-PIPERII	DINEPROPANOL HYDROCHLO
E6		LUCOSIDASE (BDELLOVIBRIO E A)/CN	BACTERIOVORUS STRAIN H
E7		LUCOSIDASE (STAPHYLOCOCCUS	S AUREUS AUREUS STRAIN
E8	1 ALPHA-D-1,4-GI MSSA476)/CN	LUCOSIDASE (STAPHYLOCOCCUS	S AUREUS AUREUS STRAIN
E9	1 ALPHA-D-1,4-GI GENE MALA)/CN	LUCOSIDASE (STAPHYLOCOCCUS	S AUREUS STRAIN ET3-1
E10	1 ALPHA-D-1,4-GI ENE MALA)/CN	LUCOSIDASE (STAPHYLOCOCCUS	S AUREUS STRAIN MU50 G
E11	ALPHA-D-1,4-GI GENE MALA)/CN	LUCOSIDASE (STAPHYLOCOCCUS	S AUREUS STRAIN RF122
E12	1 ALPHA-D-1, 4-GI TCC12228 GENE	LUCOSIDASE (STAPHYLOCOCCUS SE1191)/CN	S EPIDERMIDIS STRAIN A

 $^{=&}gt; \exp \alpha - \operatorname{cyclodext/cn}$

E1 1 A-CYCLOCOSTUNOLIDE/CN

E2	1	A-CYCLOCOSTUNOLIDE, DIHYDRO-/CN
E3	0>	A-CYCLODEXT/CN
E4	1	A-CYCLODEXTRAN DIALDEHYDE/CN
E5	1	A-CYCLODEXTRIN/CN
E6	1	A-CYCLODEXTRIN 2,3-MANNOEPOXIDE/CN
E7	1	A-CYCLODEXTRIN 6-TOSYLATE/CN
E8	1	A-CYCLODEXTRIN A, D-DIACID/CN
E9	1	A-CYCLODEXTRIN BRILLIANT YELLOW TETRAANION COMPLEX/CN
E10	1	A-CYCLODEXTRIN COMPD. WITH (±)-A-METHYLBENZEN
		EMETHANOL (1:1)/CN
E11	1	A-CYCLODEXTRIN COMPD. WITH 3-IODOPIONIC ACID (1:1)/CN
E12	1	A-CYCLODEXTRIN COMPD. WITH 4-NITROPHENYL B-D-GALA
		CTOSIDE (1:1)/CN

=> s e5

L7 1 A-CYCLODEXTRIN/CN

=> file hcaplus COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 5.61 68.30 SINCE FILE DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -12.00

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FILE COVERS 1907 - 18 Sep 2008 VOL 149 ISS 12 FILE LAST UPDATED: 16 Sep 2008 (20080916/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 13 and 17 5836 L7 L8 3 L3 AND L7

=> d 18 -13 ti abs bib

L8 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2008 ACS on STN
TI Nutraceutical compositions from microalgae and related methods of

production and administration

AB Polysaccharides with nutraceutical application may by obtained by culturing red microalgae and the nutraceutical compns. thus produced may comprise a carrier and homogenized microalgal cells. Addnl. components may include phytosterols, limonoids, flavonoids, and tocotrienols. The polysaccharides may be used in applications such as reducing cholesterol in mammals, inactivating viruses, stabilizing foods, etc. Thus, total serum cholesterol in an animal model (hamsters) over 30 days was decreased 35-62% by dietary inclusion of Porphyridium biomass homogenate and polysaccharide, the highest decreases being observed when phytosterols were also present. Transgenic algae may be used that are capable of utilizing fixed carbon sources for energy. Also provided are novel nucleic acid sequences from red microalgae.

2007:1364352 HCAPLUS <<LOGINID::20080918>> ΑN

DN 148:32596

Nutraceutical compositions from microalgae and related methods of ΤI production and administration

Dillon, Harrison F.; Somanchi, Aravind; Rao, Kamalesh; Jones, Peter J. H. ΙN

Solazyme, Inc., USA PA

SO PCT Int. Appl., 199pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 8

r An.		CENT 1	NO.			KIN		DATE		APPLICATION NO.				DATE					
ΡI	WO	2007	1364.	28				2007	1129		WO	20	07-1	US13:	 19		2	0070	119
		₩:	AE, CN, GE, KP, MK, RO, TT, AT, IS, CF,	AG, CO, GH, KR, MN, RS, TZ, BE, IT, CG,	AL, CR, GM, KZ, MW, RU, UA, BG, LT, CI,	AM, CU, GT, LA, MX, SC, UG, CH, LU,	AT, CZ, HN, LC, MY, SD, US, CY, LV, GA,	AU, DE, HR, LK, MZ, SE, UZ, CZ, MC, GN, NA,	AZ, DK, HU, LR, NA, SG, VC, DE, NL, GQ,	BA, DM, ID, LS, NG, SK, VN, DK, PL, GW,	BE DZ II LT NI SI ZF EE PT	3, 2, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,	BG, EC, IN, LU, NO, SM, ZM, ES, RO, MR,	BR, EE, IS, LV, NZ, SV, ZW FI, SE, NE,	BW, EG, JP, LY, OM, SY, FR, SI, SN,	BY, ES, KE, MA, PG, TJ, GB, SK, TD,	BZ, FI, KG, MD, PH, TM,	GB, KM, ME, PL, TN, HU, BF, BW,	GD, KN, MG, PT, TR, IE, BJ, GH,
PRAI	US U	2007 2007 2007 2007 2007 2007 2006 2006	KG, 0167 0167 0166 0166 0167 0191 -336 -336 -336 -337 -337 -816 -832 -838	KZ, 396 397 449 797 266 398 303 426 428 431 656 103 171 967P 091P 452P	MD,	RU, A1 A1 A1 A1 A1 A A A A A A P	TJ,		0719 0719 0719 0719 0719 0719 0816 0119 0119 0119 0119 0119 0119 0628 0720 0817		US US US US US	20 20 20 20 20 20	006-1 006-1 006-1 006-1	3364 3364 3364 3366 3371	28 30 31 56 03		2 2 2 2 2 2	0060 0060 0060 0060 0060 0060	119 119 119 119 119

ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2008 ACS on STN L8

TΙ Skin sanitizing compositions

AB The present invention relates to compns. and methods of sanitizing and

moisturizing skin surfaces. A sanitizing and moisturizing gel contained EtOH 55, isopropanol 3, Biowax-754 0.4, Carbopol Ultrez-10 0.3, Carbowax PEG-200 0.26, propylene glycol 0.02, aminomethylpropanol 0.15, and perfume 0.1%, and water qs.

AN 2002:551533 HCAPLUS <<LOGINID::20080918>>

DN 137:114518

TI Skin sanitizing compositions

IN Sine, Mark Richard; Wei, Karl Shiqing; Jakubovic, David Andrew; Thomas, Cheyne P.; Dodd, Michael Thomas; Putman, Christopher Dean

PA The Procter & Gamble Company, USA

SO U.S., 14 pp., Cont. of U.S. Ser. No. 321,291. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
ΡI	US 6423329	B1	20020723	US 2000-504286	20000215		
PRA	I US 1999-249717	A2	19990212				
	US 1999-120098P	P	19990216				
	US 1999-321291	A2	19990527				

RE.CNT 56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2008 ACS on STN

TI Mucosal preparation containing physiologically active peptide

AB This invention related to a mucosal preparation obtained by blending a physiol. active peptide at least with a sorbefacient and a vasodilatory compound Owing to the combined use of the sorbefacient with the vasodilatory compound, the absorption of any desired physiol. active peptide can be enhanced and thus it can be self-administered to a patient without giving any pain caused by parenteral injection. Therefore, it is highly useful as a preparation of a physiol. active peptide for prolonged administration. As the physiol. active peptide, use can be made of insulin, calcitonin, human PTH, somatostatin, glucagon, etc. As the sorbefacient, use can be made of bile acid salts, cyclodextrin, phospholipids, nonionic surfactants, higher fatty acids, etc. As the vasodilatory compds., use can be made of calcium channel inhibitors, prostaglandin E1, isosorbide nitrate, nitroglycerin,

AN 1997:259764 HCAPLUS <<LOGINID::20080918>>

DN 126:242891

OREF 126:46901a,46904a

TI Mucosal preparation containing physiologically active peptide

IN Yamamoto, Nakayuki; Ito, Teruomi

PA Asahi Kasei Kogyo Kabushiki Kaisha, Japan; Hisamitsu Seiyaku Kabushiki Kaisha; Yamamoto, Nakayuki; Ito, Teruomi

SO PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

	PATE	NT N	10.			KINI)	DATE			APPL	ICAT	ION	NO.		D_{i}	ATE		
РТ	 WO 9	7068	 ≀13			 A1	_	 1997			 WO 1	996-		 77		1	 9960:	 812	
				CN,		KR,		1337	0227		NO I		01 22	, ,			,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	012	
		RW:	AT,	BE,	CH,	DE,	DK	ES,	FΙ,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE
	JP 1	1292	2787			А		1999	1026		JP 1	995-	2080	10		19	9950	815	
	CN 1	1797	723			A		1998	0422		CN 1	996-	1928	21		19	9960	812	
	EP 8	4526	55			A1		1998	0603		EP 1	996-	9266	26		19	9960	812	
		R:	AT,	BE,	CH,	DE,	DK	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,	
			ΙE,	FI															

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JP 3824023 B2 20060920 JP 1997-509140 19960812
PRAI JP 1995-208010 A 19950815
WO 1996-JP2277 W 19960812
    MARPAT 126:242891
OS
=> s 11 or 12
         33873 L1
         41765 L2
L9
         63111 L1 OR L2
=> s 17 and 19
          5836 L7
L10
           30 L7 AND L9
=> s 110 and (PY<2003 or AY<2003 or PRY<2003)
      22958910 PY<2003
       4497131 AY<2003
       3965546 PRY<2003
L11
            20 L10 AND (PY<2003 OR AY<2003 OR PRY<2003)
=> d his 111 1-20 ti abs bib
'1-20' IS NOT VALID HERE
For an explanation, enter "HELP DISPLAY HISTORY".
=> d 111 1-20 ti abs bib
L11 ANSWER 1 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
ΤI
     Cosmetic composition comprising a complex of cyclodextrin and vitamin F
AΒ
     The invention concerns cosmetic and dermatol. compns. that contain
     complexes of vitamin F with \alpha, \beta, or \gamma-cyclodextrin.
     Addnl. substances in the formulations are: silicone oils, moisturizers,
     skin care substances, gelation agents, bactericides, antioxidants,
     sunscreens, emulsifiers, pigments, tanning agents, etc. Thus 0.1 mol
     lpha-cyclodextrin was mixed with 100 g water; 0.1 mol linolic acid was
     added, homogenized and stirred for 30 h at RT and for 8 h at 70°C;
     the product was dispersed in water, filtered, washed and dried under
     vacuum. A composition contained (weight/weight%):\alpha-cyclodextrin-linolic acid
     complex 4.0; \gamma-cyclodextrin-\alpha-tocopherol complex 1.5; octyl
     palmitate 2.5; octyl stearate 3.5; polyglycerol-2 sesquiisostearate 2.0;
     cyclomethicone, dimethiconol 3.0; lauryl dimethicone 2.0; octyl
     dimethicone ethoxy glycoside, cyclomethicone 12.0; titanium dioxide 5.0;
     polymethylsilsesquioxane 1.0; zinc oxide 2.0; glycerin 2.0; methylparaben
     0.1; sodium chloride 0.4; water 59.0.
     2004:402912 HCAPLUS <<LOGINID::20080918>>
ΑN
   140:412001
DN
TΙ
    Cosmetic composition comprising a complex of cyclodextrin and vitamin F
     Regiert, Marlies; Kupka, Michaela
ΙN
     Wacker-Chemie GmbH, Germany
PA
     Eur. Pat. Appl., 17 pp.
SO
     CODEN: EPXXDW
DT
     Patent
LA
     German
FAN.CNT 1
     PATENT NO.
                       KIND
                                DATE APPLICATION NO. DATE
                                            _____
                                _____
    EP 1419761 A1 20040519
EP 1419761 B1 20051019
PΙ
                                           EP 2003-26137
                                                                   20031113 <--
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
     DE 10253042
                        A1 20040603 DE 2002-10253042 20021114 <--
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KR 200404282	.7 A	20040520	KR 2003-77579	20031104 <
US 2004009 6 4	13 A1	20040520	US 2003-712703	20031112 <
JP 200416177	'5 A	20040610	JP 2003-385675	20031114 <
PRAI DE 2002-1025	3042 A	20021114	<	

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L11 ANSWER 2 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Production method of cyclodextrin inclusion materials using marine or animal products
- AB Title method comprise treatment of mixts. comprising lipophilic component-containing marine or animal products, starch, and lipid soluble solvents by addition of cyclodextrin synthetase. Thus, 5 g rice starch, 10 g salmon caviar, and 1 THU (based on 1 g starch) cyclodextrin synthetase were reacted in ethanol to give a cyclodextrin inclusion material showing good antioxidant property.
- AN 2004:139298 HCAPLUS <<LOGINID::20080918>>
- DN 140:182653
- TI Production method of cyclodextrin inclusion materials using marine or animal products
- IN Miwa, Shoji
- PA Ishikawa Prefecture, Japan
- SO Jpn. Kokai Tokkyo Koho, 13 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 2004051866	A	20040219	JP 2002-213621	20020723 <
PRAI	JP 2002-213621		20020723	<	

- L11 ANSWER 3 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Skin sanitizing compositions
- AB The present invention relates to compns. and methods of sanitizing and moisturizing skin surfaces. A sanitizing and moisturizing gel contained EtOH 55, isopropanol 3, Biowax-754 0.4, Carbopol Ultrez-10 0.3, Carbowax PEG-200 0.26, propylene glycol 0.02, aminomethylpropanol 0.15, and perfume 0.1%, and water qs.
- AN 2002:551533 HCAPLUS <<LOGINID::20080918>>
- DN 137:114518
- TI Skin sanitizing compositions
- IN Sine, Mark Richard; Wei, Karl Shiqing; Jakubovic, David Andrew; Thomas, Cheyne P.; Dodd, Michael Thomas; Putman, Christopher Dean
- PA The Procter & Gamble Company, USA
- SO U.S., 14 pp., Cont. of U.S. Ser. No. 321,291. CODEN: USXXAM
- DT Patent
- LA English
- FAN.CNT 2

	PA:	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΙ	US	6423329	B1	20020723	US 2000-504286	20000215 <
PRAI	US	1999-249717	A2	19990212	<	
	US	1999-12009 8 P	P	19990216	<	
	US	1999-321291	A2	19990527	<	
		F.C	FC OTEM			0.00

- RE.CNT 56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 4 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Oxidative stability and nuclear magnetic resonance analyses of linoleic

acid encapsulated in cyclodextrins

The effects of α - and β -cyclodextrin (CD) on the oxidative AB stability of linoleic acid (LA) at $35\,^{\circ}\text{C}$ were studied by measuring headspace oxygen depletion in airtight 35-mL serum bottles. LA was encapsulated in α -CD or β -CD in an aqueous solution during homogenization at 8000 rpm for 1 min and then dried under vacuum for 60 hat room temperature Headspace oxygen was measured by thermal conductivity gas chromatog. The rate of oxygen depletion for the control, which contained LA only, was 93.8 µmole/L·h. The rates of oxygen depletion for LA, encapsulated at a 1:1 mol ratio (mole CD/mol LA) in α -CD and β -CD, were 13.8 and 111 μmoles/L·h, resp. When LA was encapsulated in $\alpha\text{-CD}$ and $\beta\text{-CD}$ at a 2:1 mol ratio (moles CD/mol LA), the rates of oxygen depletion were 0.573 and 53.9 μ moles/L·h, resp. Although α -CD protected LA from reaction with oxygen at both ratios, the rate of oxygen depletion by LA encapsulated in β -CD at a 1:1 mol ratio was not statistically different from the control. $\beta\text{-CD}$ protected LA from reaction with oxygen at a 2:1 mol ratio. 1H NMR spectra of the complexes formed from 1:1 mol ratios of LA and CD indicated that LA was encapsulated in α -CD or β -CD.

AN 1997:681639 HCAPLUS <<LOGINID::20080918>>

DN 127:358219

OREF 127:70123a,70126a

- TI Oxidative stability and nuclear magnetic resonance analyses of linoleic acid encapsulated in cyclodextrins
- AU Reichenbach, Wendy A.; Min, David B.
- CS Department of Food Science, The Ohio State University, Columbus, OH, 43210, USA
- SO Journal of the American Oil Chemists' Society (1997), 74(10), 1329-1333
 CODEN: JAOCA7; ISSN: 0003-021X
- PB AOCS Press
- DT Journal
- LA English
- RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 5 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Mucosal preparation containing physiologically active peptide
- AB This invention related to a mucosal preparation obtained by blending a physiol. active peptide at least with a sorbefacient and a vasodilatory compound Owing to the combined use of the sorbefacient with the vasodilatory compound, the absorption of any desired physiol. active peptide can be enhanced and thus it can be self-administered to a patient without giving any pain caused by parenteral injection. Therefore, it is highly useful as a preparation of a physiol. active peptide for prolonged administration. As the physiol. active peptide, use can be made of insulin, calcitonin, human PTH, somatostatin, glucagon, etc. As the sorbefacient, use can be made of bile acid salts, cyclodextrin, phospholipids, nonionic surfactants, higher fatty acids, etc. As the vasodilatory compds., use can be made of calcium channel inhibitors, prostaglandin E1, isosorbide nitrate, nitroglycerin, etc.
- AN 1997:259764 HCAPLUS <<LOGINID::20080918>>
- DN 126:242891
- OREF 126:46901a,46904a
- TI Mucosal preparation containing physiologically active peptide
- IN Yamamoto, Nakayuki; Ito, Teruomi
- PA Asahi Kasei Kogyo Kabushiki Kaisha, Japan; Hisamitsu Seiyaku Kabushiki Kaisha; Yamamoto, Nakayuki; Ito, Teruomi
- SO PCT Int. Appl., 48 pp. CODEN: PIXXD2

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Patent
DT
LA Japanese
FAN.CNT 1
                 A1 19970227 WO 1996-JP2277 19960812 <--
     PATENT NO.
     _____
     WO 9706813
PΙ
         W: CA, CN, JP, KR, US
         RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
     JP 11292787 A 19991026 JP 1995-208010 19950815 <--
     CN 1179723 A 19980422 CN 1996-192821
EP 845265 A1 19980603 EP 1996-926626
                                                                     19960812 <--
                                                                     19960812 <--
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
                          B2 20060920
                                             JP 1997-509140
     JP 3824023
                                                                     19960812 <--
PRAI JP 1995-208010
WO 1996-JP2277
                         A 19950815 <--
W 19960812 <--
OS
     MARPAT 126:242891
L11 ANSWER 6 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
     A method of producing a taxane-type diterpene
TΙ
AΒ
     A simple method of producing a taxane-type diterpene by plant tissue
     culture is disclosed. Productivity can be improved by carrying out the
     culture in the presence of coronatines, a bacterium that produced the
     coronatines, a culture solution or a culture extract of such bacteria, cyclic
     polysaccharides, fatty acids, or an amino or imino derivative of jasmonic
     acids.
     1996:572123 HCAPLUS <<LOGINID::20080918>>
ΑN
     125:219760
OREF 125:41103a,41106a
TI A method of producing a taxane-type diterpene
    Yukimune, Yukihito; Hara, Yasuhiro; Tan, Hiroaki; Tomino, Ikuo
ΤN
PA Mitsui Petrochemical Industries, Ltd., Japan
SO Eur. Pat. Appl., 32 pp.
     CODEN: EPXXDW
DΤ
     Patent
LA
     English
FAN.CNT 3
     PATENT NO. KIND DATE APPLICATION NO. DATE
     PATENT NO.
     EP 727492
                         A2 19960821
PΙ
                                             EP 1995-308498
                                                                    19951127 <--
     EP 727492
                         A3
                                19961016
                  B1
     EP 727492
                                20010131
        R: DE, FR, GB, IT, NL
     JP 08140690 A 19960604
JP 3549594 B2 20040804
                                             JP 1994-291783
                               19960604
                                                                      19941125 <--
JP 3549594

JP 08163991

A 19960625

JP 1994-312258

19941215 <--

JP 3625908

JP 08205882

A 19960813

JP 1995-301654

JP 3746550

B2 20060215

PRAI JP 1994-291783

A 19941125 <--

JP 1994-301179

A 19941205 <--

JP 1994-312258

A 19941215 <--

JP 1995-218874

A 19950828 <--

MARPAT 125:219760
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- L11 ANSWER 7 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Aggregation of polyunsaturated fatty acids in the presence of cyclodextrins
- AB The aggregation behavior of the polyunsatd. fatty acids (PUFA) linoleic acid and arachidonic acid was studied in the presence of cyclodextrins

(CDs). The influence of CD concentration on CMC of PUFA suggests that two CD mols. bind sequentially to one mol. of PUFA. Two equilibrium consts., K1 representing the interaction of the first CD mol., and K2, the interaction of the second, were determined by non-linear regression of the PUFA CMC vs. CD concentration data to an expression deduced from the reaction scheme in the equilibrium The effect of pH and the structure of the CD on the equilibrium consts.

was studied. It is postulated that the first CD mol. interacts with the carboxyl group of PUFA through hydrogen bonding when the fatty acid is protonated, while the second CD mol. binds to the hydrocarbon chain of the PUFA through hydrophobic interaction. The formation of hydrogen bonds was principally affected by the inner diameter of the CD, while the hydrophobic interactions were very strongly affected by the polarity of the CD group coating the inner channel. The relevance of the results for the development of enzyme assays involving fatty acids is discussed.

- 1995:628687 HCAPLUS <<LOGINID::20080918>> AN
- 123:50376 DN
- OREF 123:8923a,8926a
- Aggregation of polyunsaturated fatty acids in the presence of cyclodextrins
- ΑU
- Bru, Roque; Lopez-Nicolas, Jose M.; Garcia-Carmona, Francisco Dep. Bioquim. Biol. Mol. "A", Univ. Murcia, Murcia, E-30001, Spain CS
- Colloids and Surfaces, A: Physicochemical and Engineering Aspects (SO 1995), 97(3), 263-9 CODEN: CPEAEH; ISSN: 0927-7757
- Elsevier PΒ
- Journal DT
- LAEnglish
- L11 ANSWER 8 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- Entrapment of liquid lipids into powdery matrixes of saccharides and ΤТ proteins
- The emulsifying activity, the high stabilizing activity of the emulsion AΒ and the formation of a fine dense skin layer during drying were the properties of agents that effectively entrapped liquid lipids. Gum arabic and gelatin were effective. Addition of an agent having a property to a base agent lacking the property improved the entrapment. Oxidation of entrapped liquid lipid was retarded. However, the extent of retardation depended on the kind of lipids and the kind of entrapping agents. Oxidation processes of some combinations of lipids and entrapping agents were expressed by a kinetic model including oxygen diffusion through dehydrated entrapping agents. Et eicosapentaenoate was also stabilized by the entrapment.
- ΑN 1995:485889 HCAPLUS <<LOGINID::20080918>>
- DN 122:263834
- OREF 122:48177a,48180a
- Entrapment of liquid lipids into powdery matrixes of saccharides and ΤI proteins
- Matsuno, Ryuichi; Imagi, Jun; Adachi, Shuji ΑU
- Fac. Agric., Kyoto Univ., Kyoto, 606-01, Japan CS
- Dev. Food Eng., Proc. Int. Congr. Eng. Food, 6th (1994), Meeting SO Date 1993, Volume Pt. 2, 1065-7. Editor(s): Yano, Toshimasa; Matsuno, Ruuichi; Nakamura, Kozo. Publisher: Blackie, Glasgow, UK. CODEN: 61FFAL
- $\mathsf{D}\mathsf{T}$ Conference
- English LA
- ANSWER 9 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN L11
- ΤI Utilization of cyclodextrin as fat soluble compound carrier to serum-free culture of rat astrocytes
- AΒ α -Cyclodextrin complexes with fat-soluble vitamins and unsatd. fatty acids were prepared and examined as replacements for bovine serum albumin as

fat-soluble compound carriers on cultured rat astrocytes. In serum-supplemented medium, it was difficult to evaluate the effects of fat-soluble compds. in serum on cell growth. Therefore, serum-free chemical defined medium supplemented with growth factors, hormones, and nutrients was developed for rat astrocytes to evaluate these effects. $\alpha\text{-Cyclodextrin}$ complexes with 3 vitamins (vitamin A acetate, E, and K1) and 3 fatty acids (linoleic, linolenic, and oleic acids) showed growth promoting activities for astrocytes in serum-free medium. Usually, supplementing fat-soluble compds. to a cell culture medium is very difficult, especially to a low or no protein medium, but $\alpha\text{-cyclodextrin}$ can replace albumin as a fat-soluble compound carrier in serum-free cell cultures.

AN 1993:579303 HCAPLUS <<LOGINID::20080918>>

DN 119:179303

OREF 119:32055a,32058a

- TI Utilization of cyclodextrin as fat soluble compound carrier to serum-free culture of rat astrocytes
- AU Nakama, Akihiko
- CS Osaka City Inst. Public Health Environ. Sci., Osaka, 543, Japan
- SO Annual Report of Osaka City Institute of Public Health and Environmental Sciences (1992), Volume Date 1991, 54, 48-53 CODEN: AOISDR; ISSN: 0285-5801
- DT Journal
- LA Japanese
- L11 ANSWER 10 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Retarded oxidation of liquid lipids entrapped in matrixes of saccharides or proteins
- AΒ Me linoleate (ML), linoleic acid (LA), and Et eicosapentaenoate (EE) were entrapped in saccharide and protein matrixes, and then stored at 37° in a desiccator controlled at 75% relative humidity. ML entrapped with α -cyclodextrin, maltodextrin, and pullulan was extremely resistant to autoxidn., but LA entrapped with maltodextrin and pullulan rapidly oxidized. LA entrapped with α -cyclodextrin was the most stable against oxidation ML entrapped with gelatin or gum arabic was less resistant to autoxidn. than that entrapped with pullulan; there was little difference in the susceptibility to oxidation between ML and LA entrapped with gelatin or gum arabic. Egg albumin protected ML more effectively against oxidation than LA, while sodium caseinate protected LA more than ML. EE entrapped with pullulan was highly resistant to oxidation, 90% of the total lipid remaining after 35 days. The effect on the oxidation of diffusion of oxygen through the matrix was estimated Retardation of oxidation

of the entrapped lipid can not be explained only by the effect of diffusion.

- AN 1992:590442 HCAPLUS <<LOGINID::20080918>>
- DN 117:190442
- OREF 117:32869a,32872a
- TI Retarded oxidation of liquid lipids entrapped in matrixes of saccharides or proteins
- AU Imagi, Jun; Muraya, Koji; Yamashita, Daisuke; Adachi, Shuji; Matsuno, Ryuichi
- CS Fac. Agric., Kyoto Univ., Kyoto, 606-01, Japan
- SO Bioscience, Biotechnology, and Biochemistry (1992), 56(8), 1236-40
 CODEN: BBBIEJ; ISSN: 0916-8451
- DT Journal
- LA English
- L11 ANSWER 11 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Powderization of liquid-state lipids
- AB Liquid-state lipids (linoleic acid, Me linoleate, or Me oleate) were

powderized by adsorption on gum arabic, starch, maltodextrin, $\alpha\text{-cyclodextrin}$, maltose, glucose, or CM-cellulose. Lipids adsorbed on $\alpha\text{-cyclodextrin}$, gum arabic, or CM-cellulose had high stability. The emulsifying activity of the lipid-adsorbent complex is described.

AN 1991:654556 HCAPLUS <<LOGINID::20080918>>

DN 115:254556

OREF 115:43273a,43276a

TI Powderization of liquid-state lipids

AU Matsuno, Ryoichi; Imagi, Jun

CS Agric. Coll., Kyoto Univ., Kyoto, Japan

SO New Food Industry (1991), 33(5), 57-64

CODEN: NYFIAM; ISSN: 0547-0277

DT Journal

LA Japanese

L11 ANSWER 12 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN

TI Specific adsorbents in isolation and purification of cyclodextrins

AB A number of synthesized affinity sorbents were tested to find methods for the separation of α -, β -, and γ -cyclodextrins (CDs) from one another and from acyclic dextrins. None of the gels retarded acyclic dextrins, whereas α -CD was specifically adsorbed onto supports derivatized with alkyl functions, β -CD was specifically adsorbed onto supports derivatized with phenyl or substituted Ph, and γ -CD was specifically adsorbed onto a gel derivatized with a naphthyl compound It was evident that for achievement of binding capacities high enough for practical preparation of the CDs, various parameters such as the support material, its porosity, ligand, ligand concentration, temperature, and the

composition of the mobile phase must be optimized.

AN 1989:453519 HCAPLUS <<LOGINID::20080918>>

DN 111:53519

OREF 111:9029a,9032a

TI Specific adsorbents in isolation and purification of cyclodextrins

AU Makela, Mauri; Mattsson, Pekka; Korpela, Timo

CS Dep. Biochem., Univ. Turku, Turku, SF-20500, Finland

SO Biotechnology and Applied Biochemistry (1989), 11(2), 193-200 CODEN: BABIEC; ISSN: 0885-4513

DT Journal

LA English

L11 ANSWER 13 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN

TI Pharmaceuticals containing unsaturated fatty acids and stimulators for synthesis of prostaglandin and hydroxy fatty acids

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AB The title composition contains ≥ 1 unsat. C18-22 fatty acid derivs. containing 3-5 isolated double bonds and which may be Me or Et substituted at

the 2, 3, 16-20 position, selected from the free terminal carboxylic acids, amides, or CO2X derivs. (X = protecting group removable under acidic conditions, 1- or 2-lysophospholipid, metal cation, amine cation, cationic ion-exchanger). It also contains a stimulator with simultaneously stabilizing properties selected from ≥1 phenols I (R1 = OH, CO2H, CH2CO2H, CH:CHCO2H, CH2CHR4R5, CH(OH)CH2NHR6; R2, R3 = H, OH; R4 = H, CO2H; R5 = H, NH2; R6 = H, Me, Et]; indoles II (R7 = H, CO2H; R8 = H, NH2; R9 = H, OH); cysteine, homocysteine, or liponic acid wherein the alicyclic alkyl residue may be shortened by <4 CH2-groups; a peptide containing ≤10 amino acids and in which ≥1 may be replaced by any of the above compds.; one of the above amino compds. substituted by C1-4 alkyl; a flavonoid substituted by ≥1 OH linked to a sugar residue; a salt of the above named compds.; as ester containing an alkoxy-containing residue, or its amide, mono- or dialkylamide. Addnl., it contains stabilizers selected from DMSO, EtOH, polyols, polyol esters, phospholipids, sugar lipids, cyclodextrins, proteins, cytochrome c derivs., or E-vitamins in solid or liquid form. A mixture containing 0.3 mL 0.03M

K phosphate buffer, 0.5 mg enzyme (from sheep sperm vesicles or homogenate of kidney medulla), 2.75 μg 14C-arachidonic acid, and 0.5 mg I [R1 = CH2CH(NH2)CO2H, R2 = R3 = H] (stimulator) was incubated for 10 min at 37° and quenched with citric acid. The formation of total prostaglandin increased 5.5-fold over the amount formed in the absence of a stimulator; the relative amts. of PGE2, PGF2 α , and PGD2 with stimulator were 81, 2, and 17%, resp., and 83, 2, and 15%, resp., in the absence of a stimulator.

AN 1988:443459 HCAPLUS <<LOGINID::20080918>>

DN 109:43459

OREF 109:7237a,7240a

TI Pharmaceuticals containing unsaturated fatty acids and stimulators for synthesis of prostaglandin and hydroxy fatty acids

IN Weithmann, Klaus Ulrich

PA Hoechst A.-G., Fed. Rep. Ger.

SO Eur. Pat. Appl., 35 pp.

CODEN: EPXXDW

DT Patent

LA German FAN.CNT 1

	PAT	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	ΕP	244832	A2 A3 B1	19871111 19891129 19920624	EP 1987-106520	19870506 <
		R: AT, BE, CH,	DE, ES	, FR, GB,	GR, IT, LI, LU, NL, SE	
	DE	3615710	A1	19871126	DE 1986-3615710	19860509 <
	ΑT	77549	T	19920715	AT 1987-106520	19870506 <
	ES	2051705	T3	19940701	ES 1987-106520	19870506 <
	DK	8702356	A	19871110	DK 1987-2356	19870508 <
	DK	167518	B1	19931115		
	-	8772641	A	19871112	AU 1987-72641	19870508 <
		603574	B2	19901122		
	_	62267222	A	19871119	JP 1987-110953	19870508 <
		8703299	A	19871230		
		44433	A2	19880328	HU 1987-2088	19870508 <
		201671	В	19901228		
		1302266	С	19920602		19870508 <
		82459	A	19940731		19870508 <
		5043328	A	19910827		19890201 <
PRAI		1986-3615710	A	19860509		
		1987-106520	A	19870506		
	US	1987-46650	В3	19870507	<	

- L11 ANSWER 14 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI The effect of bovine serum albumin on the synthesis of prostaglandin and incorporation of [3H]acetate into platelet-activating factor
- AB As determined by RIA, bovine serum albumin (BSA) inhibited bradykinin (BK) (5 ng/mL)- and ionophore A 23187 (10 μ M)-stimulated synthesis of prostaglandins (PGs) by human embryo lung fibroblasts (IMR-90) in a concentration-dependent manner. Addition of [3H]arachidonate followed by

TLC showed that, in the presence of 2 mg/mL BSA, IMR-90 cells released essentially only fatty acids following stimulation with bradykinin. Little if any prostaglandin and no endoperoxide were detected. In the absence of BSA, .apprx.70% of the released label was detected as prostaglandin. $\alpha\text{-Cyclodextrin}$, another trapper of fatty acid, inhibited PG synthesis in much the same way. BSA and $\alpha\text{-cyclodextrin}$ also inhibited prostacyclin synthesis in endothelial cells derived from the calf pulmonary artery. However, the inhibition of PG synthesis in these cells was not as complete as that in the IMR-90 cells. In contrast to the effect of the trappers on PG synthesis, BSA and $\alpha\text{-cyclodextrin}$ potentiated BK- and ionophore-stimulated incorporation of [3H]acetate into PAF in the endothelial cells. The labeled PAF was not released from the cells in either the presence or absence of the trappers, which suggests that BSA causes an increase in acetate-labeled cellular PAF by trapping released fatty acid.

AN 1987:569447 HCAPLUS <<LOGINID::20080918>>

DN 107:169447

OREF 107:27070h,27071a

- TI The effect of bovine serum albumin on the synthesis of prostaglandin and incorporation of [3H]acetate into platelet-activating factor
- AU Heinsohn, Carlotta; Polgar, Peter; Fishman, Jordan; Taylor, Linda
- CS Sch. Med., Boston Univ., Boston, MA, 02118, USA
- SO Archives of Biochemistry and Biophysics (1987), 257(2), 251-8 CODEN: ABBIA4; ISSN: 0003-9861
- DT Journal
- LA English
- L11 ANSWER 15 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Stabilization of lipids by molecular inclusion: cyclodextrins and casein as antioxidants
- AB The effects of cyclodextrin and casein inclusion on the kinetics of linoleic acid [60-33-3] and arachidonic acid [506-32-1]
-] oxidation in dispersions containing lipoxygenase or Na bisulfite were evaluated

by monitoring free radical side reactions and O consumption. The fatty acid peroxidn. inhibition by casein was primarily by reversible inclusion of the free polyunsatd. fatty acid. Cyclodextrins and casein inhibited both enzymic and nonenzymic peroxidn. Inhibitor consts. were relatively high unless the concentration of fatty acids was limiting.

AN 1986:477674 HCAPLUS <<LOGINID::20080918>>

DN 105:77674

OREF 105:12597a,12600a

- TI Stabilization of lipids by molecular inclusion: cyclodextrins and casein as antioxidants
- AU Laakso, Simo
- CS Dep. Biochem., Univ. Turku, Turku, 20500, Finland
- SO Lipid Oxid.: Biol. Food Chem. Aspects, Contrib. LIPIDFORUM/SIK Symp. (1986), Meeting Date 1985, 165-70. Editor(s): Marcuse, Reinhard. Publisher: Scand. Forum Lipid Res. Technol., Goeteborg, Swed. CODEN: 55ATAL
- DT Conference

- LA English
- L11 ANSWER 16 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Growth of an established line of mouse mammary tumor cells under serum-free conditions
- An established line of mouse mammary tumor cells (MTD cells) were cultured AB in a serum-free medium consisting of a 1:1 mixture of Dulbecco's modified Eagle's medium and Ham's F-12 medium supplemented with bovine serum albumin (BSA), insulin, and transferrin. To promote cell attachment and spreading, culture dishes were precoated with plasma fibronectin isolated from fibrinogen. Under these serum-free conditions, MTD cells grew at a rate close to that attained by the serum-supplemented medium. Among the additives in the serum-free medium, BSA was replaced with oleic acid or a complex of oleic acid and α -cyclodextrin. Transferrin was replaced with Fe2+ or Fe3+. Addition of polyvinylpyrrolidone further improved the growth. Thus, MTD cells can be grown on a fibronectin-coated surface in a chemical defined medium with insulin as the only protein supplement. MTD cells grown under the serum-free conditions still retained the differentiated properties of the original MTD cells; i.e., the production of mouse mammary tumor virus in response to dexamethasone.
- AN 1986:164689 HCAPLUS <<LOGINID::20080918>>
- DN 104:164689
- OREF 104:25993a,25996a
- TI Growth of an established line of mouse mammary tumor cells under serum-free conditions
- AU Kawamura, Kazuo; Enami, Jumpei; Kohmoto, Kaoru; Koga, Mutuyosi
- CS Sch. Med., Dokkyo Univ., Mibu, 321-02, Japan
- SO Dokkyo Journal of Medical Sciences (1985), 12(2), 167-80 CODEN: DJMSDB; ISSN: 0385-5023
- DT Journal
- LA English
- L11 ANSWER 17 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Inhibition of lipid peroxidation by casein. Evidence of molecular encapsulation of 1,4-pentadiene fatty acids
- AB The capability of cyclodextrins to form mol. inclusion complexes with linoleate resulted in inhibition of oxygenation in a lipoxygenaselinoleate model reaction. The inhibited rates were established instantaneously upon addition of the complexant and were maintained until linoleate was exhausted. Total cessation of the reaction was not obtained with cyclodextrins. Casein-inhibited reaction mixts. also exhibited these characteristics. Both casein and cyclodextrins protected linoleate against autoxidn., although they did not change free radical generation by xanthine oxidase or Fe2+ reactions. Since neither of the inhibitors affected the enzyme directly, casein may act, in analogy with cyclodextrins, by forming linoleate complexes which reduce the oxidizable monomer fatty acids via a standing equilibrium and thus result in substrate limitation of reaction rates. Comparisons of lipid peroxidn. at acidic and alkaline pH, in the presence of increasing amts. of the complexants, detergent, and hydroperoxides, supported this view.
- AN 1984:81327 HCAPLUS <<LOGINID::20080918>>
- DN 100:81327
- OREF 100:12263a,12266a
- TI Inhibition of lipid peroxidation by casein. Evidence of molecular encapsulation of 1,4-pentadiene fatty acids
- AU Laakso, Simo
- CS Dep. Biochem., Univ. Turku, Turku, SF-20500/50, Finland
- SO Biochimica et Biophysica Acta, Lipids and Lipid Metabolism (1984), 792(1), 11-15
- CODEN: BBLLA6; ISSN: 0005-2760
- DT Journal

- LA English
- L11 ANSWER 18 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI α -Cyclodextrin: a partial substitute for bovine serum albumin in serum-free culture of mammalian cells
- AB The use was investigated of oleic acid- or linoleic acid- α -cyclodextrin inclusion complexes as albumin substitutes for mammalian cells. α -Cyclodextrin did not show any cytotoxic effects at 2g/L medium. Growth curves are shown for 2 types of cells. UMCL cells grew well enough in the cyclodextrin-complex-containing, serum-free medium, whereas HEL cells required a small amount of albumin in addition to cyclodextrin for abundant growth.
- AN 1982:612006 HCAPLUS <<LOGINID::20080918>>
- DN 97:212006
- OREF 97:35533a,35536a
- TI $\alpha\text{-Cyclodextrin:}$ a partial substitute for bovine serum albumin in serum-free culture of mammalian cells
- AU Yamane, Isao; Kan, M.; Minamoto, Y.; Amatsuji, Y.
- CS Inst. Tuberculosis Cancer, Tohoku Univ., Sendai, 980, Japan
- SO Cold Spring Harbor Conferences on Cell Proliferation (1982), 9 (Growth Cells Horm. Defined Media, Book A), 87-92 CODEN: CSHCAL; ISSN: 0097-5230
- DT Journal
- LA English
- L11 ANSWER 19 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI α -Cyclodextrin, a novel substitute for bovine albumin in serum-free culture of mammalian cells
- AB The use of α -, β -, and γ -cyclodextrin (CD) in combination with unsatd. fatty acids as a serum substitute in mammalian cell cultures was examined by using a human lymphoblast cell line (UMCL) grown in RITC 56-1 medium supplemented with synthetic lecithin, cholesterol, galactose, and mannose and by using human diploid fibroblasts (HEL) grown in RITC 80-7 medium. On the basis of cytotoxic and cost considerations, α -CD was used for the expts. Both α -CD-oleic acid and α -CD-linoleic acid had growth-enhancing effects on UMCL cells up to 100 mg/L medium but exhibited toxic effects at higher concns. However, when 100 mg α -CD included with both fatty acids and 1000 mg free α -CD were added to 1 L of medium, stable and reproducible growth-promoting effects were observed. With HEL cells, growth similar to that in bovine serum albumin-supplemented medium was observed by addition of a concentrated α -CD complex to a final concentration of 10-20 mg/L.
- AN 1982:100488 HCAPLUS <<LOGINID::20080918>>
- DN 96:100488
- OREF 96:16453a,16456a
- TI α -Cyclodextrin, a novel substitute for bovine albumin in serum-free culture of mammalian cells
- AU Yamane, Isao; Kan, Mikio; Minamoto, Yoshiki; Amatsuji, Yasuo
- CS Res. Inst. Tuberc. Cancer, Tohoku Univ., Sendai, 980, Japan
- SO Proceedings of the Japan Academy, Series B: Physical and Biological Sciences (1981), 57(10), 385-9
 CODEN: PJABDW; ISSN: 0386-2208
- DT Journal
- LA English
- L11 ANSWER 20 OF 20 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Stabilization of autoxidizable materials by means of inclusion
- AB Adducts of α -dextrin (cyclohexaamylose) (I), β -dextrin (cycloheptaamylose) (II) and deoxycholic acid (III) were prepared with linoleic acid (IV), linolenic acid (V), Me linolenate (VI), PhCH:CHCHO (VII), and vitamin A palmitate (VIII). They were found to be very

resistant to autoxidation. The conventional procedure of preparing choleic acids yielded stable products with V and VIII. The products obtained from dextrins with IV, V, and VII needed purification. A heat treatment under high vacuum was found to be reliable for obtaining stable adducts free of oxidizable contamination. The principle of inclusion stabilization appears to be established by these examples and by the previous work on fatty acid stabilization by means of urea (C.A. 44, 11123f). II (8 g.) in 100 cc. O-free 50% aqueous EtOH treated at about 70° with 1.3 q. IV, the mixture stirred 4 hrs. at room temperature and centrifuged, and the solid dried over P205 at 0.5 mm. gave 7.7 g. II-IV adduct containing 7.28 g. IV (titrated in hot 50% aqueous EtOH with 0.05N KOH and phenolphthalein. II-IV adduct sublimed after rinsing with N under a high vacuum 9 hrs. at 120-5° gave 6.9% IV. Purified II-IV adduct (1.63 g.) in 100 cc. hot 50% aqueous EtOH extracted twice with 50-cc. portions trimethylpentane, the extract dried and evaporated, the residual oil brominated in Skellysolve F, and the resulting white crystals (75 mg.) repptd. from warm Et20 with Skellysolve F yielded 47 g. tetrabromostearic acid, m. 115-16.5°. II (1.6 g.) and 0.32 g. V treated in the usual manner in 20 cc. aqueous EtOH, the solids isolated and heated 17 hrs. at 122 $^{\circ}$ and 0.5 mm. pressure, two 0.7-g. portions of the residue (each containing 67 mg. V) exposed to pure O in a Warburg apparatus (the manometers being filled with silicone fluid) at $37 \pm 0.2^{\circ}$ (one in a dry and one in a humid atmospheric) and the charge brominated in the usual manner gave eventually hexabromostearic acid. The II-VI adduct containing 10.8% VI was obtained in the same manner. II (5.0 g.) in 100 cc. H2O and 0.9 g. VII shaken 16 hrs. at room temperature, the solids isolated in the usual manner and heated 3 hrs. at $100-40^{\circ}$ and 0.5 mm. gave an adduct containing 10.5% (9.6%) VII (determined as the 2,4-dinitrophenylhydrazone, m. 258-9°) and 0.3% (1.3%) PhCH:CHCO2H. I (2.0 g.) in 15 cc. O-free H2O warmed to 70° with IV in 15 cc. EtOH, the mixture kept 4 hrs. at room temperature, the crystals

isolated by centrifugation and dried, and a part heated to $130-60^{\circ}$ during 16 hrs. at 0.5 mm. gave I-IV adduct (115 μ l. O uptake during 40 hrs. under standard conditions); another part of the crude product digested with 10 cc. EtOH gave I-IV adduct (760 μ l. O-uptake). III (6.0 g.) in 20 cc. absolute EtOH and 0.55 g. V in 5 cc. EtOH kept 16 hrs. at -5 to -10° gave III-V adduct containing 8.3% V. The adduct was refluxed 1 hr. with 8 times its weight of xylene, the III-xylene adduct filtered and washed with C6H6, the combined xylene and C6H6 solution evaporated,

the oily residue extracted with Skellysolve C, the extract evaporated, and the residue titrated with alkali to determine the acid content. III (1.0 g.) and 0.1 g. VIII in 4 cc. hot EtOH cooled to room temperature, held 12 hrs. at -3° , and the light yellow crystals filtered and dried in a high vacuum gave the III-VIII adduct containing 10.8% VIII.

AN 1956:23995 HCAPLUS <<LOGINID::20080918>>

DN 50:23995

OREF 50:4858g-i,4859a-e

- TI Stabilization of autoxidizable materials by means of inclusion
- AU Schlenk, Hermann; Sand, Donald M.; Tillotson, Jerry Ann
- CS Univ. of Minnesota, Austin
- SO Journal of the American Chemical Society (1955), 77, 3587-90 CODEN: JACSAT; ISSN: 0002-7863
- DT Journal
- LA Unavailable